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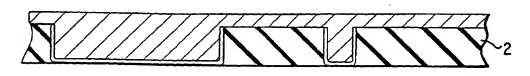
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(54) Title: MODIFIED PLATING SOLUTION FOR PLATING AND PLANARIZATION AND PROCESS UTILIZING SAME



(57) Abstract: A modified plating solution that can be used to electroplate a high quality conductive material that can be effectively polished and planarized includes (1) a solvent, (2) an ionic species of the conductive material to be deposited, (3) at least one additive to improve electrical and structural properties, and (4) a modifying agent.



MODIFIED PLATING SOLUTION FOR PLATING AND PLANARIZATION AND PROCESS UTILIZING SAME

This application claims the priority of provisional application serial no. 60/182,100, filed February 11, 2000, the entire disclosure of which is incorporated by reference herein.

Reference is hereby made to related application serial nos. 09/472,523, filed December 27, 1999, entitled WORK PIECE 10 CARRIER HEAD FOR PLATING AND POLISHING, and 09/511,278, filed February 23, 2000, entitled PAD DESIGNS AND STRUCTURES FOR A VERSATILE MATERIALS PROCESSING APPARATUS.

BACKGROUND AND SUMMARY OF INVENTION

Multi-level integrated circuit manufacturing requires many steps for metal and insulator film depositions followed by photoresist patterning and etching or other means of material removal. After photolithography and etching, the resulting wafer or substrate surface is non-planar and contains many features such as vias, lines, or channels. Often these features need to be filled with a specific material such as a metal, and then the wafer topographic surface needs to be planarized again, making it ready for the next level of processing.

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Electrodeposition is a widely accepted technique for the deposition of a highly conductive material such as copper (Cu) into the features on the semiconductor wafer surface. Chemical mechanical polishing (CMP) is then employed to planarize the resulting surface.

In Figure 1a, the large feature 1 and the small feature 1s are opened in the insulator layer 2, which is grown on a wafer. To fill these features with Cu, a barrier or adhesive layer 3 is first deposited over the whole wafer surface. Then a conductive Cu seed layer 4 is deposited over the barrier

layer 3. Cu is electrodeposited over the whole surface by (1) making an electrical contact to the barrier layer 3 and/or the Cu seed layer 4; (2) placing the wafer in a standard Cu plating electrolyte; (3) placing an anode in the electrolyte; and (4) applying a negative voltage to the Cu seed layer with respect to the anode.

Figure 1b shows the wafer after a short period of time which is adequate to deposit a Cu layer 5 with the thickness 10 5a. As shown in Figure 1b, the Cu layer of nominal thickness 5a is adequate to fill in the small feature 1s since there is Cu film growth even on the conductive vertical walls of this feature. The large feature 1, however, is still not filled with Cu. To fill the large feature 1, Cu plating needs to 15 proceed further, eventually yielding the structure depicted in Figure 1c.

As can be seen in Figure 1c, in this conventional approach, the electrodeposited Cu layer 5 forms a very large 20 metal overburden 6 on the top surface of the insulator 2 and over the small feature 1s. The overburden 6a over the large feature 1 is very small. The surface of the structure in Figure 1c is non-planar, and therefore needs to be polished and planarized. The overburden and portions of the barrier layer 3 are customarily removed by CMP, yielding the structure in Figure 1d, which has electrically isolated Cu-filled features. Removal of the large and non-uniform metal overburden of Figure 1c from the wafer surface is time consuming and expensive and is a major source of dishing defects 6b in large features.

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It would be highly desirable if the plating process could yield a Cu film which was planar and had a uniform overburden as depicted in Figure 1e. CMP of such a substrate would be much faster and more economical and defects would be minimized. If the plating process could yield Cu-filled features with no

overburden as depicted in Figure 1f, then there would not be the need for CMP of the Cu layer. Only the portions of the barrier layer 3 on the top surface of the insulator 2 would have to be removed.

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Electrodeposition is commonly performed in specially formulated plating solutions or electrolytes containing ionic species of Cu as well as additives that control the texture, morphology, and the plating behavior of the Cu layer. A proper electrical contact is made to the seed layer on the wafer surface, typically along its circumference, and the wafer surface is dipped in the plating solution. A consumable Cu anode or an inert anode plate is also placed in the electrolyte. Deposition of Cu on the wafer surface can then be initiated when a cathodic potential is applied to the wafer surface with respect to the anode (i.e., when a negative voltage is applied to the wafer surface with respect to the anode plate).

There are many Cu plating solution formulations, some of which are commercially available. One such formulation uses Cu-sulfate (CuSO₄) as the copper source. James Kelly et al., J. Electrochemical Society, vol.146, p. 2540-45 (1999). A typical Cu-sulfate plating solution contains water; Cu-sulfate; sulfuric acid (H₂SO₄); a small amount of chloride ions; and a carrier, such as polyethylene glycols and/or polypropylene glycols. Some other chemicals are then added to this solution in small amounts to achieve certain properties of the Cu deposit. These additives can be classified under general categories such as levelers, brighteners, grain refiners, wetting agents, stress-reducing agents, and the like.

Commonly used levelers and brighteners are generally sulfur-containing compounds, such as derivatives of thiourea.

Other levelers and brighteners are sulfonic acid derivatives,

such as mercaptobenzene sulfonate. Other brighteners include 2,4-imidazolidine-diol, thiohydantoin, polyethers, polysulfides, and various dyes. There is a large volume of literature on the additives for Cu-plating solutions and their influence on the electroplated deposits. For example, U.S. 4,430,173 discloses an additive composition Patent No. sodium comprising the salt of ω-sulfo-n-propyl diethyldithiocarbamate and crystal violet, which shows excellent stability. U.S. Patent No. 4,948,474 discloses a brightener additive for a Cu plating solution. U.S. Patent No. 4,975,159 discloses lists of alkoxylated lactams and sulfurcontaining compounds which were found to be effective U.S. Patent No. 3,328,273 describes Cu plating additives. baths containing organic sulfide compounds.

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Although a large volume of literature exists on the subject of additives to Cu plating solutions, many of the additive formulations are kept as trade secrets by plating solution suppliers. Some examples of Cu plating additive solutions provided commercially are: (1) CUBATH M® system, marketed by Enthone-OMI; (2) COPPER GLEAM® system, marketed by LeaRonal; and (3) ULTRAFILL® Addition agent and Suppressor, marketed by Shipley. Commercially available Cu plating solutions with additives typically yield bright and soft Cu deposits that have low stress. Copper layers deposited out of 25 these solutions cannot be polished and planarized with the same solution, simply because the plating solutions are formulated only for plating, not for polishing or planarization.

30 Copper layers are traditionally polished and planarized by CMP in a machine specifically designed for polishing. In this method, the plated wafer is loaded onto a carrier head. The wafer surface covered with the non-planar Cu deposit (Figure 1c) is brought into contact with a polishing pad and 35 a polishing slurry. The polishing slurry contains oxidizing

chemicals and micron or sub-micron size abrasive particles. When the pad and the wafer surfaces are pressed together and moved with respect to each other, polishing by the abrasive particles is initiated and the metal overburden is removed from the surface. A different CMP slurry is used to remove the barrier layer from the top surface of the insulator. The desired planar surface with electrically isolated Cu-filled features shown in Figure 1d is eventually obtained.

10 The chemistry of the polishing slurry and the type of the abrasive particles used in a given CMP process are selected according to the chemical nature of the material to be removed. Therefore, the compositions of the polishing slurries for copper, tungsten, tantalum, tantalum nitride, silicon dioxide, and like materials that are used in integrated circuit (IC) 15 manufacturing may all be different. For example, U.S. Patents Nos. 4,954,142; 5,084,071; 5,354,490; 5,770,095; 5,773,364; 5,840,629; 5,858,813; 5,897,375; 5,922,091; and 5,954,997, all various CMP slurry compositions for effective 20 polishing of Cu. Slurries typically contain a solvent and a selection of abrasive particles, such as silica or alumina particles, which are suspended in the solvent. complexing agents such as NH3 and/or oxidizing agents such as $\mathrm{NO_3}^-$ and $\mathrm{Fe}\left(\mathrm{CN}\right)_6{}^{3-}$ are also included in the formulations to 25 increase the dissolution rate of the abraded material and thus increase the Cu removal rate.

A typical CMP slurry has a high pH so that a passivating surface layer can be formed and sustained on Cu surfaces in the features where the pad cannot make high pressure contact. The surface layer over the Cu film protects Cu in such areas from chemical attack by the solution. High regions of the Cu layer making high pressure physical contact with the pad get polished because the abrasive particles can remove the passivating

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surface layer. The abraded material is then carried away from the wafer surface and dissolved by the slurry.

discussed As in J. Steigerwald et al., 5 Microelectronic Materials, sections 7.2.1. and 7.2.2., John Wiley & Sons Inc. (1997), Cu is not expected to form a protective surface film in acidic solutions with low pH. Therefore, if an acidic slurry is employed for CMP, Cu in all regions, including in the recessed features, would dissolve into the acidic slurry and planarization as depicted in Figure le would not be possible. In CMP of Microelectronic Materials, it is suggested that when using an acidic CMP slurry, a nonnative, film-forming agent such as benzotriazole (BTA) may be added to the slurry composition to avoid chemical etching of the deposit in the recessed areas. However, the data in CMP of Microelectronic Materials also demonstrates that BTA may reduce the polishing rate.

According to CMP of Microelectronic Materials, during CMP,

20 a protective layer such as an oxide layer first forms on the

Cu surface due to the chemical composition and the pH of the

slurry. The surface film is then efficiently removed by the

mechanical action of the abrasive particles. Removed material

is moved away from the vicinity of the wafer surface to avoid

25 re-deposition. This process continues until all the metal on

high surfaces making contact with the pad is removed. CMP

slurries that are commercially available are all designed for

polishing and planarization only. There is no CMP slurry

formulation that allows plating of materials.

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Thus, the chemical compositions of metal plating solutions and metal polishing and planarization slurries are very different. Polishing cannot be realized in prior art metal plating solutions, and plating cannot be achieved with standard CMP slurries. This is not customarily a problem since metal

plating and CMP processes are carried out in different machines at different times, and the plating solutions and CMP slurries used in these machines are only expected to achieve their respective single functions, namely, deposition and polishing.

5 However, this conventional approach is time consuming and it raises the manufacturing cost for integrated circuits.

It is an object of the present invention to provide a metal deposition solution that allows for polishing and planarizing the plated metal layer using the same solution. Metal layers deposited and planarized using such a solution would yield desirable flat surfaces as shown in Figure 1e in a short period of time and would even achieve the structure depicted in Figure 1f using a single machine. The idea of simultaneous plating and polishing or plating/polishing of a conducting material on a workpiece or wafer surface was disclosed in co-pending U.S. Patent Application Serial No. 09/201,929, filed on December 1, 1998, the entirety of which is incorporated herein by reference.

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6,004,880 discloses a modified CMP U.S. Patent No. apparatus and technique to achieve simultaneous plating and polishing on a semiconductor wafer. However, compositions of solutions that can be successfully employed in such processes 25 have not been fully disclosed. U.S. Patent No. 6,004,880 discloses a modified CMP apparatus and a simultaneous plating/polishing technique that would be achieved by modifying a CMP slurry. In other words, the approach is modifying a CMP process to also do plating. Therefore, an electrolyte composition that might contain Cu-sulfate was proposed to be 30 mixed into a CMP slurry. Such an approach of mixing plating electrolytes into polishing slurries has several drawbacks.

Electronic applications of electroplated metals, such as electroplated Cu, require not only planar surfaces, but also

excellent electrical and mechanical properties. Copper films used in such applications should have low resistivity values close to the bulk resistivity of Cu, which is about 1.6x10-6 should also have good electromigration Films 5 properties; should adhere well to their substrates; and should have large grains. Although planarization is important in the overall process, a highly planar Cu layer with high resistivity would not be usable in many integrated circuit applications where high performance is desired, because the limiting speed at which the circuit can be run is a strong function of the resistivity of the metal used in its structure. Copper is typically plated out of acidic electrolytes because layers obtained from these electrolytes have low resistivity and other desirable characteristics. Plating Cu out of a solution with large amounts of undesirable impurities typically deteriorate the properties of the deposited films. Therefore, the quality of Cu layers obtained from slurry/electrolyte mixtures would in all probability be poor.

20 Mixing plating solutions into CMP slurries to modify the slurries for plating would be also problematic for other The chemistry of plating solutions and polishing slurries are highly incompatible. The Cu-sulfate plating solutions that are commonly used for Cu plating are highly acidic solutions with pH values well below 0.5, typically below High pH values deteriorate the plated film properties, typically giving rise to rough and/or burned deposits, especially at high plating current densities that are necessary for fast processing. This deterioration is because limiting 30 current density values decrease as the pH of the solution goes In contrast, CMP slurries that are commonly used have pH values well above 2, typically above 4.0. For example, CMP slurries CPS-01 and CPS-03 sold by 3M® corporation have pH values of around 7. HASTILITE® marketed by Rhodes has a pH of 7.25. MICROPLANAR CMP 9000® by EKC Technology Inc. has a pH

value of 8.83. As discussed in CMP of Microelectronic Materials, high pH is desirable in CMP slurries because it allows the formation of a protective surface layer on Cu. In acidic electrolytes with low pH values, surface layers such as 5 Cu-oxides cannot be stable. It is therefore expected that mixing low pH plating solutions with high pH polishing slurries would have detrimental effects on both plating and polishing processes.

- 10 Slurries are formulated to keep their abrasive particles dispersed or in suspension. According to the basic theory of dispersions, dispersion characteristics of small particles in a solution such as a CMP slurry is a strong function of the pH of the solution. Changing the pH of a perfect dispersion may totally destroy that dispersion and cause the particles to 15 agglomerate and precipitate out. Plating is also sensitive to pH changes and is affected by the electrolyte composition. Even parts per million (ppm) levels of impurities/additives in plating solutions have profound effects on the properties of 20 the plated materials. Electronic applications of conductive materials such as Cu require deposition of good quality films with low resistivity and large grain. Chemicals in the polishing slurry would affect the quality of the plating solution and therefore the quality of the Cu layer that might 25 be plated with an electrolyte/slurry mixture.
- U.S. Patent No. 6,004,880 discloses the idea of adding a plating solution to modify a polishing slurry for the simultaneous plating/polishing of Cu on a wafer with surface 30 feature widths of preferably 1 micron or less. In many circuit designs, however, there are surface features with widely varying widths. The small feature in Figure 1a, for example, may have a width of only 0.1-0.5 microns, whereas the width of the large feature may be 10-100 microns. The depth of the features may 35 be in the range of microns. 1-5 Α

slurry/electrolyte mixture could be introduced onto a wafer surface with only submicron size features, if the particle size in the slurry is selected such that the particles are larger than the feature size. This way, the large particles cannot get into the wells or channels defined by the small features. However, if there are both large and small features on the wafer, then use of such a slurry would result in lodging of some of the abrasive particles into the large vias and channels. The lodged particles would then interfere with Cu plating into the features; increase resistivity; destroy microstructure and device performance; and drastically reduce the process yield.

Even if the plating solution and the CMP slurry could be mixed and used for plating and polishing, their recycling would be uneconomical. Once used, these solutions would have to be discarded and the cost of processing would be high.

The present invention solves these problems by modifying 20 a plating solution for plating and planarization, rather than modifying a polishing slurry for polishing and plating as disclosed in the prior art.

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BRIEF DESCRIPTION OF THE DRAWINGS

Figures 1a through 1d show an example of a prior art procedure for filling wafer surface features with electrodeposited Cu, and then polishing the wafer to obtain a structure with a planar surface and electrically isolated Cu plugs or wires;

Figure 1e shows a metal deposit with uniform metal overburden across the surface of the substrate according to the present invention;

Figure 1f shows plating just in the holes according to the present invention; and

Figure 2 schematically shows a plating/polishing apparatus that can be used according to the present invention.

DETAILED DESCRIPTION OF THE DRAWINGS

The present invention modifies a plating solution for plating and planarization due to the following factors.

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The quality of the deposited metal layer is of prime importance. In any process, one has to first assure this quality. Therefore, instead of taking a polishing solution and adapting it for plating, we took the route of taking a well-15 formulated plating solution and modifying it by adding a modifying agent to achieve in-situ planarization of the plated The conditions to be satisfied by the modifying agent are: (1) addition of the modifying agents into the plating solution should not affect the grain size and resistivity of the plated layer in a detrimental way; (2) the modified plating solution containing modifying agents should yield a plated layer that can be efficiently polished and planarized; and (3) a user should be able to easily monitor the amount of the modifying agents in the solution and re-furbish and recycle the used solution in a closed loop system.

The present invention discloses plating solution compositions that are modified to allow the deposition of a high quality Cu layer, and at the same time allow either 30 simultaneous or sequential polishing and planarization of the deposited layer. In this approach, commercially available, highly acidic Cu plating solutions are modified by the addition of oxidizers which do not appreciably affect the pH of the solution or the quality of the plated Cu layer. No slurry or

particles are included in the formulation. Polishing and planarization is achieved using a fixed abrasive pad.

A version of a plating/polishing apparatus that can be used to practice the present invention is schematically shown The carrier head 10 holds the wafer 16 and at the in Fig. 2. same time provides an electrical contact 7 to the seed layer on the wafer surface. The head can be rotated around a first axes 10b. It can also be moved in x, y, and z directions. A fixed abrasive polishing pad 8 is placed on an anode assembly 10 9 across from the wafer surface. Modified plating solution 9a of the present is supplied to the wafer surface, preferably through the openings in the anode assembly and the pad as shown by the arrows in the figure. The solution then flows over the edges of the pad into the chamber 9c to be re-circulated after 15 cleaning/filtering/refurbishing. An electrical contact 9d is provided to the anode assembly 9. The anode assembly 9 can also be rotated around the axes 10c. The gap between the wafer surface and the pad is adjustable by moving the carrier head and/or the anode assembly in the z direction. When the wafer surface and the pad are touching, the pressure that is exerted on the two surfaces can also be adjusted.

For plating, a potential is applied between the electrical contact 7 to the wafer 16 and the electrical contact 9d to the anode assembly 9, making the wafer surface more negative than the anode assembly. Under applied potential, a high quality layer of metal plates out of the modified plating solution onto the wafer surface. By adjusting the gap between the pad and the wafer surface and/or by adjusting the pressure with which the pad and the wafer surface touch each other, one can achieve just plating, or plating and polishing. For example, if there is a gap between the wafer surface and the pad, plating is expected to take place over the whole wafer surface as illustrated in Figure 1c. In this case, a metal film is

obtained that can be polished in a CMP process in a separate CMP machine. It should be noted that Cu layers plated out of the modified plating solutions of the present invention were found to be polished more efficiently compared to Cu layers obtained from standard plating solutions and therefore are advantageous.

If the pad and the wafer surface in Figure 2 were touching at low pressures, then plating can freely take place in the holes in the substrate where there is no physical contact between the wafer surface and the abrasive pad, but the plating rate will be reduced on the top surfaces where there is physical contact between the pad and the surface. The result is a metal deposit with uniform metal overburden across the surface of the substrate as shown in Figure 1e. This is in contrast to the conventional deposition method of Figure 1c where there is significant variation in metal overburden across the substrate. If the pressure with which the substrate and the pad surfaces touch each other is further increased, it is 20 possible to obtain plating just in the holes as shown in Figure In this case, the increased polishing action on the high points of the substrate surface does not allow accumulation of metal layer on these regions.

It should be understood that the modified plating solution of the present invention may be used in apparatus with various other designs, including designs for just plating and designs for plating and polishing. For plating a metallic layer that can be later polished more effectively, many kinds of plating apparatus can employ the solution of this invention by simply replacing the standard plating solution with the modified solution of the present invention. For plating/polishing applications, any apparatus that can apply a voltage difference between the wafer surface and an electrode touching the modified plating solution of the present invention, while

pressing a fixed abrasive pad against the wafer surface and moving the pad and the wafer surface with respect to each other, can be employed.

It is not fully understood how the addition of small amounts of oxidizers in the highly acidic Cu plating solutions allows the use of these solutions for plating and planarization. However, it is possible that the surface layer formed on the Cu deposit by the presence of oxidizers does not interfere with the plating of a good quality Cu layer, but at the same time can be efficiently removed from the sections of the film where the pad contacts it with some pressure.

The amount of oxidizer added to the plating solution may be less than 500 ppm, however preferably it should be more. Oxidizer concentration may typically be in the 0.01 wt.% to 10 wt.% range. Both inorganic and organic oxidizers, either pure or mixed, or their mixtures can be used as modifying agents, but organic oxidizers are preferred. Among the many organic 20 oxidizers known to those in the field of chemistry, the preferred modifying agents are organic nitrites and nitrates. Although butyl nitrite is an organic oxidizer that was used as the modifying agent in the following examples to demonstrate the present invention, other modifying agents can also be used 25 to obtain the same result. For example, other organic oxidizers, preferably organic nitrites can be used. nitrites include, but are not limited to, alkyl nitrites, aromatic nitrites, and polyaromatic nitrites. Alkyl nitrites include, but are not limited to, primary, secondary and tertiary compounds of methyl, ethyl, propyl, butyl, and amyl nitrites. Additionally, nitrates of the above compounds may also be used.

Although the examples use Cu deposits, it should be understood that many other conductive materials such as Cu

alloys, W, Au, Ni, Pt, Pd, Ag, Co, Sn, Pb and their alloys can be used in the practice of the present invention.

5 EXAMPLES

EXAMPLE 1: STANDARD PLATING SOLUTION

A Cu-sulfate based Cu plating solution was prepared as follows:

70 grams per liter of CuSO₄+5H₂O, 150 grams per liter of concentrated H₂SO₄, and 70 ppm per liter of Cl⁻ ions were mixed in enough water to make 10 liters of solution. Twenty-five ml of Ultrafill S2001®, 1.0 ml of Ultrafill A2001® from Shipley were then added to obtain a standard good quality plating electrolyte.

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This solution was used for Cu plating on a 200 mm diameter wafer surface using the apparatus of Figure 2. surface contained sub-micron size features as well as features in the 10-100 micron range. The pad was a fixed abrasive pad supplied by 3M® company. The diameter of the pad was 180 mm 20 and the anode assembly was oscillated in the horizontal direction so that plating could be achieved on all areas on the larger wafer surface. During plating, the distance between the pad and the wafer surface was kept at around 0.1 cm. 25 plating current was 2 Amp and the plating solution flow was 5 liters/minute. The wafer was rotated at 75 rpm and the anode assembly with the pad was rotated at 100 rpm in the same direction. Several wafers were plated for times ranging from 90 seconds to 4 minutes. Resulting Cu deposits were similar to the ones depicted in Figures 1b and 1c depending on the 30 depth of the surface features and the plating period. deposits after aging at room temperature for one day had a resistivity of below 2x10-6 ohm-cm, indicating good material quality.

EXAMPLE 2: POLISHING AND PLANARIZATION USING STANDARD PLATING SOLUTION

The plating experiment of Example 1 was repeated, except this time, after an initial period of 30 seconds, the pad was pushed against the wafer surface at a pressure of 1 psi for plating as well as polishing and planarization. The resulting Cu deposit had a rough surface with deep scratches apparently caused by the abrasive pad. There were also Cu particles smeared all over the surface of the wafer. Very little amount of material removal was achieved because material removed from one region of the surface by the action of the abrasive pad was probably deposited back onto the surface at another region in the form of smeared particles, which were welded or bonded to the substrate surface. The substrate defect level was extremely high and feature filling was poor.

EXAMPLE 3: MODIFIED PLATING SOLUTION ACCORDING TO THE PRESENT INVENTION

Five ml per liter of butyl-nitrite was added as a modifying agent to the electrolyte of Example 1 and the plating and polishing experiment of Example 2 was repeated using this modified plating solution. The resulting Cu deposit was highly planar and was similar to the structure shown in Figure 1e. Copper layer resistivity was still below 2x10⁻⁶ ohm-cm, demonstrating the ability of the modified electrolyte to yield high quality Cu deposits. Copper film was planar with uniform overburden over the sub-micron size features as well as the large features.

The foregoing disclosure has been set forth merely to illustrate the invention and is not intended to be limiting. Since modifications of the disclosed embodiments incorporating the spirit and substance of the invention may occur to persons skilled in the art, the invention should be construed to include everything within the scope of the appended claims and equivalents thereof.

WHAT IS CLAIMED IS:

1. A process of depositing and simultaneously polishing and planarizing a high quality conductive layer on a surface of a substrate, comprising:

loading a substrate on a holder;

introducing a plating solution comprising an oxidizer on an abrasive polishing pad;

pressing the abrasive polishing pad against the surface 10 of said substrate;

contacting the plating solution with the surface of the substrate and a second electrode;

applying a potential difference between the surface of the substrate and the second electrode;

depositing a conductive layer on the surface of the substrate; and

moving the abrasive pad and the surface of the substrate with respect to each other, thereby simultaneously polishing and planarizing the conductive layer on said substrate.

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- 2. A process according to Claim 1, wherein the conductive layer comprises copper.
- 3. A process according to Claim 1, wherein the plating 25 solution comprises an acidic copper plating solution.
 - 4. A process according to Claim 3, wherein the acidic copper plating solution has a pH value of less than 4.
- 5. A process according to Claim 1, wherein said oxidizer is selected from the group consisting of an inorganic oxidizer, an organic oxidizer, and mixtures thereof.
- 6. A process according to Claim 5, wherein the oxidizer 35 is an organic nitrite.

7. A process according to Claim 6, wherein the nitrite is selected from the group consisting of alkyl nitrites, aromatic nitrites, and polyaromatic nitrites.

- 8. A process according to Claim 7, wherein the organic nitrite is an alkyl nitrite.
 - 9. A process according to Claim 8, wherein the alkyl nitrite is butyl nitrite.

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- 10. A process according to Claim 1, wherein the oxidizer is an organic nitrate.
- 11. A process according to Claim 1, wherein the substrate 15 comprises surface features having a width of about 0.1-100 microns.
 - 12. A modified plating solution for simultaneous polishing and planarization of a substrate, comprising:
- 20 a solvent;

an ionic species of a conductive material; and an oxidizer.

- 13. A modified plating solution according to Claim 12, wherein said oxidizer is selected from the group consisting of an inorganic oxidizer, an organic oxidizer, and mixtures thereof.
- 14. A modified plating solution according to Claim 12, 30 wherein said oxidizer is an organic nitrite selected from the group consisting of alkyl nitrites, aromatic nitrites, and polyaromatic nitrites.
- 15. A modified plating solution according to Claim 12, 35 wherein said solution has a pH value of less than 4.

16. A modified plating solution according to Claim 12, wherein said oxidizer is present in an amount of more than 500 ppm.

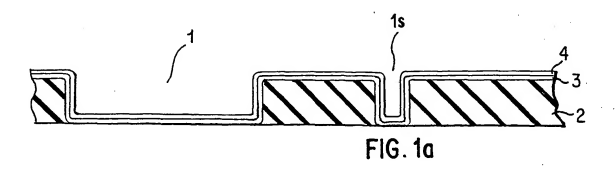
- 5 17. A modified plating solution according to Claim 12, wherein said oxidizer is present in an amount of 0.01 to 10 wt.% of said solution.
- 18. A modified plating solution according to Claim 12, 10 wherein said conductive metal is Cu.
- 19. A modified plating solution according to Claim 12, wherein said conductive metal is selected from the group consisting of W, Au, Ni, Pt, Pd, Ag, Co, Sn, Pb and their alloys.
- 20. A modified plating solution according to Claim 12, further comprising at least one additive selected from the group consisting of levelers, brighteners, grain refiners, 20 wetting agents, and stress-reducing agents.
 - 21. A plating solution for plating a conductive layer on a surface of a substrate, comprising:
 - a solvent;
- an ionic species of a conductive material; and an oxidizer.
- 22. A plating solution according to Claim 21, wherein said oxidizer is selected from the group consisting of an inorganic oxidizer, an organic oxidizer, and mixtures thereof.
 - 23. A plating solution according to Claim 21, wherein said oxidizer is an organic nitrite selected from the group consisting of alkyl nitrites, aromatic nitrites, and polyaromatic nitrites.

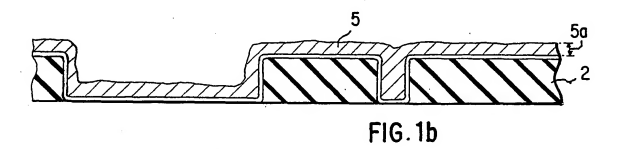
24. A plating solution according to Claim 21, wherein said solution has a pH value of less than 4.

- 25. A plating solution according to Claim 21, wherein 5 said oxidizer is present in an amount of more than 500 ppm.
 - 26. A plating solution according to Claim 21, wherein said oxidizer is present in an amount of 0.01 to 10 wt.% of said solution.

10

27. A plating solution according to Claim 21, wherein said conductive metal is Cu.





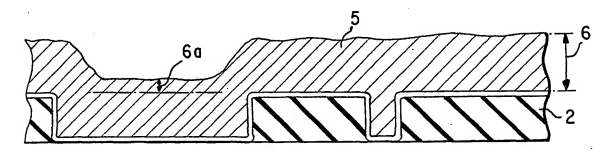


FIG. 1c

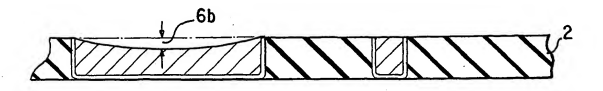


FIG. 1d

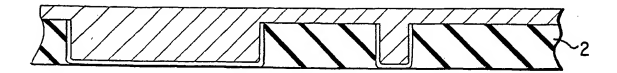


FIG.1e

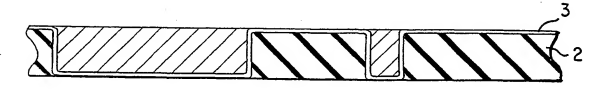


FIG. 1f

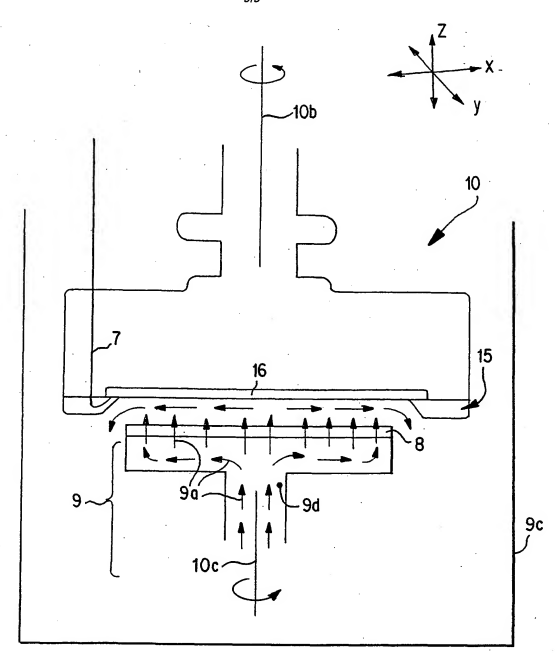


FIG. 2

INTERNATIONAL SEARCH REPORT

Form PCT/ISA/210 (second sheet) (July 1998)

International application No.

PCT/US00/31283

	PC1/U300/31263
A. CLASSIFICATION OF SUBJECT MATTER	
IPC(7) : B24B 1/00	
US CL: 451/54; 51/307 According to International Patent Classification (IPC) or to both national classification and IPC	
B. FIELDS SEARCHED	
Minimum documentation searched (classification system followed by classification symbols) U.S.: 51/307; 216/88,89; 438/691,692,693; 451/36/.41.63	
0.0 31/307, 210/08,03, 436/031,032,033, 431/30/,41,03	
Documentation searched other than minimum documentation to the extent that such documents are included in the fields searched	
·	
Electronic data base consulted during the international search (name of data base and, where practicable, search terms used)	
C. DOCUMENTS CONSIDERED TO BE RELEVANT	
Category * Citation of document, with indication, where	appropriate, of the relevant passages Relevant to claim No.
A US 5,807,165 A (UZOH et al) 15 September 1998	(15.09.1998) 1-27
A US 5,846,398 A (CARPIO) 08 December 1998 (08	.12.1998)
A US 5,866,031 A (CARPIO et al) 02 February 1999	
A US 6,004,880 A (LIU et al) 21 December 1999 (21	12,
A, E US 6,010,964 A (GLASS) 04 January 2000 (04.01.	,

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A, P US 6,103,096 A (DATTA et al) 15 August 2000 (1	1 '
A, P US 6,132,586 A (ADAMS et al) 17 October 2000 (1
A, E US 6,156,661 A (SMALL) 05 December 2000 (05.	12.2000) 1-27
	1
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	Date of mailing of the international search report
10 January 2001 (10.01.2001)	15FEB 2001
Name and mailing address of the ISA/US Commissioner of Patents and Trademarks	Authorized officer
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Washington, D.C. 20231 Facsimile No. (703)305-3230	Telephone No. 703-308-1148 Technology Center 3700